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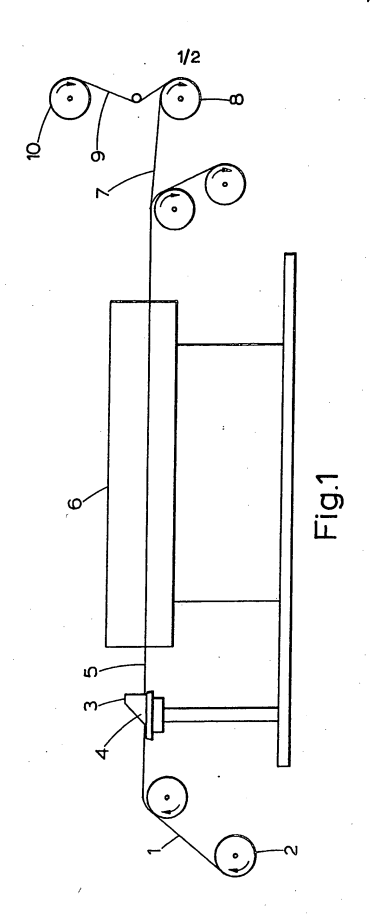
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  - GB 1142323
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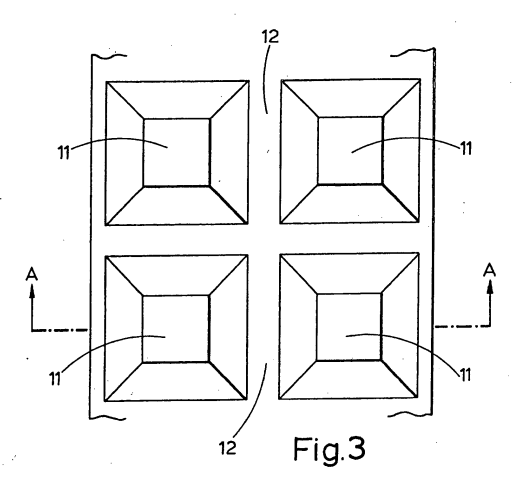
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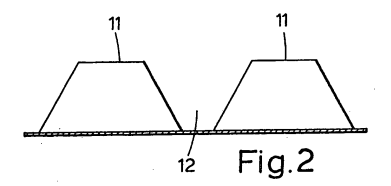
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## (54) Wound dressings for burns

(57) Low adherency wound dressings comprise a wound facing layer, an intermediate absorbent layer and an outer layer, the wound facing layer comprising a conformable elastomeric apertured film, the intermediate absorbent layer comprising a conformable hydrophilic foam and the outer layer comprising a continuous moisture vapour transmitting conformable film; materials therefor and processes for the manufacture thereof are described.







## **SPECIFICATION**

## W und dressing, manufacture and use

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	5	The present invention relates to an absorptive wound dressing suitable for use on burns or other wounds which dressing has a reduced tendency to adhere to the wound and can act as a bacterial barrier. The present invention also relates to the manufacture and use of such	5	
	10	dressings.  Burns and other related wounds such as donor sites and the like present a serious problem in that they tend to produce large amounts of exudate which can cause conventional dressings to become saturated or to stick to the wound or even become infected. One method of covering such wounds has been to cover the wound with a material into which new epithelial or fibroblast growth can penetrate. Dressings of this kind are disclosed in U.S. Patents Nos.	10	
	15	3526224, 3648692 and 3949742.  However such dressings can be extremely painful to remove and often require surgical excision. A fundamentally different approach requiring a fundamentally different type of dressing excision.	15	
	20	United States Patents Nos. 3543750, 2923298 and British Patent No. 778813 which later patents cover successfully used materials such as Melolin ("Melolin" is a registered Trade Mark of T J. Smith and Nephew Limited, Welwyn Garden City, Herts., U.K.). One more recent attempt at non-adherent dressings is United States Patent No. 3709221 which discloses a attempt at non-adherent dressings is United States Patent No. 3709221 which discloses a	20	
	25	fibrous layer and an absorbent intermediate layer which was also envisaged as normally being fibrous. In order to reduce the tendency of this material to adhere to the wound the inner layer had to be treated with an agent to render it non-wetted by body liquid. It is now realised that it had to be treated with an agent to render it non-wetted by body liquid. It is now realised that it	25	
	30	treatment. As it will become apparent hereinanter it has now been discovered that by avoiding fibrous materials it is possible to produce a dressing with reduced tendency to adhere to wounds without the need for special treatments. An attempt at producing an absorbent dressing is described in US. Patent No. 3888748 which describes a dressing fabricated from at least four	30	•
	35	grid and collects any loose thread or particle that may be present in the core material. It is now realised that it is desirable to avoid the use of wound facing layers that can allow such penetration of the central layer to the wound surface. It has also been realised that it would be penetration of the central layer to the wound surface.	35	
	40	Nos. 3709221 and 3888248 disclose materials which are bonded along their edges which may reflect a desire to improve conformability. The dressing of the present invention allows for bonding over the whole of the operative area while retaining flexibility.	40	
	45	Accordingly the present invention provides a low authority which wound dressing a wound facing layer, an intermediate absorbent layer and an outer layer which wound dressing is characterised in that the wound facing layer comprises a conformable elastomeric apertured film, the intermediate absorbent layer comprises a conformable hydrophilic foam and the outer layer comprises a continuous moisture vapour transmitting conformable film.  Normally the three layers of the dressing of this invention are attached in a contiguous and co-extensive manner; that is the dressing is normally provided in the form of a laminate.	45	
	50	The conformable elastomeric apertured film of the dressing of this invention acts as a low adherency wound facing layer. This layer allows wound exudate to pass to the absorbent layer adherence would surface.	50	
•	55	Preferably the elastomeric apertured film is sufficiently continuate to allow the trouble dressing to conform to the body contours and thereby maintain overall contact with the wound surface to ensure that exudate from the wound is absorbed.  It is also desirable that the elastomeric apertured film should be sufficiently elastically extensible to adjust to any dimensional changes in the absorbent layer which may occur, for	55	
	60	Normally the elastomeric apertured film is made of a pharmaceutically acceptable water insoluble polymer. Pr ferred polymers for use are elastomers. Suitable elastomers include polyurethan s, polybutadiene and the like. The preferred marterials for the apertured films are thermoplastic polyurethanes and polybutadienes.	60	
	65	Preferred thermoplastic polyurethanes are linear polyur thanks softiaming polyurethanes are disclosed in U.S. Patent Specification No. 2,871,218. Suitable linear polyether polyurethanes are disclosed in U.S. Patent Specification No. 2,899,411. Favoured thermoplastic polyurethanes include Estanes from B.F.	65	

G drich Corp. Preferred solution casting grades are Estane 5714F1, 5702, and 5703. A pref rred extrusion grade is Estane 580201. Suitable polybutadienes are 1,2 polybutadienes. Favoured 1,2 polybutadienes contain a major amount of syndiotactic 1,2 polybutadiene, have a crystallinity of 25% to 30% and an average molecular weight in excess of 100,000. Preferred 1,2 polybutadienes are known as RB 810, 5 RB820 and RB 830 made by Japan Synthetic Rubber Co. The number and size of the apertures in the apertured film will be sufficient to allow the wound exudate to pass through the film to the absorbent layer. Most aptly the apertured film is adapted so that the size of apertures in combination with the thickness of the film prevent the absorbent layer contacting the wound 10 surface. Suitable apertured films have apertures with a dimension of from 0.05 to 4 mm, more 10 aptly from 0.05 to 2.5 or 0.05 to 2 mm and preferably from 0.1 to 2.5mm. Suitable apertured films have a thickness of 0.01 to 2.5mm, typically 0.01 to 0.25mm and preferably of 0.05 to 0.5mm. Favoured apertured films of the invention have 4 to 40 apertures per cm with a dimension of 0.05mm to 2.5mm. The wound face of the apertured film suitably will have 15 to 15 80% of its area void (the apertures), more suitably will have 25 to 75% of its area void and 15 most suitably will have 35 to 65% of its area void. In a favoured aspect of the invention the elastomeric apertured film is in the form of a net which is preferably an integral net. The term 'integral net' means a net in which the strands and junctures are formed integrally during manufacture. The integral net of the wound dressing of the invention can have any convenient form 20 depending on the chosen arrangement of strand, juncture and hole areas and also their shapes and relative size. In one preferred form the net consists essentially of longitudinal and transverse strands intersecting at right angles to give a square grid hole pattern. Suitable nets of this type aptly have 2 to 40 strands per cm desirably 4 to 40 strands per cm 25 and preferably 2 to 24 strands per cm in both longitudinal and transverse directions. 25 Variations on the square grid pattern can give other desirable forms of the integral net. Unequal density of strands in either the longitudinal or transverse directions will give rectangular hole areas. Continuous parallel strands in one direction with a staggered arrangement of connecting strands in the other direction will give a "brickwork" pattern. Other apt forms of the 30 integral polymer nets can have strands at an angle to the longitudinal or transverse direction 30 (that is diagonal strands). Another preferred form of the integral polymer net can have a staggered arrangement of circular or approximately circular (for example hexagonal) arrangements of strands and hole areas. The integral polymer net can be in the form of a mixed pattern of two or more of the arrangements if desired. The apertured film used in this invention aptly will have a weight of 10 gsm to 80 gsm and 35 preferably will have a weight of 15 gsm to 50 gsm. The desirable conformability of the wound dressing of the invention is consistent with the use of elastomeric materials such as integral nets of polyurethane or other elastomer. Suitable integral nets of polyurethane or other elastomer will have an elongation at break of 40 100% to 800%, desirably of 200% to 750% and preferably of 300% to 700% when 40 measured as a 2.5 cm wide strip at 30 cm/min strain rate at 20°C. Other suitable conformable apertured films are thin flexible elastomeric films which have been perforated. The continuous moisture vapour transmitting conformable film outer layer of the wound 45 dressing of the invention may be used to regulate the moisture loss from the wound area under 45 the dressing and also to act as a barrier to bacteria so that bacteria on the outside surface of the dressing cannot penetrate to the wound area. Suitable continuous conformable films will have a moisture vapour transmission rate of 300 to 5000 grams preferably 500 to 2000 grams/square metre/24 hrs at at 37.5°C at 100% to 50 10% relative humidity difference. It has been found that such moisture vapour transmission 50 rates of the continuous film allow the wound under the dressing to heal under moist conditions without causing the skin surrounding the wound to macerate. This outer layer will be made of polymer. Suitable polymers for use in the outer layer include urethanes and copolymers of alkoxy alkyl 55 acrylates or methacrylates such as those disclosed in British Patent No. 1,280,631. 55 Preferably the outer layer is a conformable polyurethane film. Preferred polyurethane films are made from linear polyurethanes as hereinbefore described with respect to the wound facing layer. Favoured continuous films will b 12.5 micron to 37.5 micron thick. A preferred polyurethane for use in such thickn ss is Estane 5714F. A 25 micron 60 thick film of Estane 5714F has a moisture vapour transmission rate of approximately 1800 60 g/m²/24 hours/37.5°C at 100% to 10% relative humidity difference so that it may be employed to produce a moisture vapour transmission within the preferred range. The outer layer can be a conformable polyurethane-incompatible polymer blend film continu-

Suitable conformable polyurethane blend films are disclosed in United Kingdom patent

	5	application No. 8122250.  Apt conformable polyurethane blend film out r layers have a thickness of 0.0125 m to 0.125 mm. Such films can have a moisture vapour transmission rate of at least 500 g and preferably at least 1000g/m²/24 hours/at 37.5°C at 100% to 10% relative humidity difference.  A preferred polyurethane blend film comprises a blend of a linear polyurethane (60 parts by weight of Estane 580201 available from B. F. Goodrich) and a high inact polystyrene (40 parts	5
	10	by weight of compound ref. 6 mw available from R. H. Cole Limited). A favoured fifth of this composition has a thickness of 0.084 mm and a moisture vapour transmision rate of 1660 g/m²/24 hours/at 37.5°C at a 100% to 10% relative humidity difference.  The continuous moisture vapour transmitting conformable film outer layer can comprise a	10
		moisture vapour transmitting adhesive layer.  In such adhesive containing outer layers the adhesive layer will be covered by an extensible moisture vapour transmitting layer to provide a non adhesive surface on the outer layer of the	
•	15	dressing.  At least one of the adhesive or extensible layers will be continuous.  Preferred adhesive layers are continuous. Suitable adhesives which are moisture vapour transmitting as a continuous layer include various acrylate ester copolymers, polyvinyl ethyl ether and polyurethane pressure sensitive adhesives. Examples of suitable pressure sensitive	15
:	20	adhesives are given in British Patent No. 1,280,631.  A preferred pressure sensitive adhesive comprises a blend of a high and low viscosity polyvinyl ethyl ethers in particular 'adhesive composition A' disclosed in British Patent polyvinyl ethyl 1,280,631. Other preferred pressure sensitive adhesives comprise copolymers	20
:	25	of acrylate ester with acrylic acid for example as disclosed in United Kingdom Application No. 8106707 and in particular a copolymer of 47 parts by weight of butylacrylate, 47 parts by weight of 2 ethyl hexyl acrylate and 6 parts by weight of acrylic acid with an intrinsic viscosity of at least 1.9 dl/g polymerised in acetone according to the general method given in above	25
	20	United Kingdom application.  The suitable continuous adhesive layers can have a weight per square metre of 15 g to 70 g and preferably of 20 g to 40 g.  The extensible moisture vapour transmitting layer which covers the adhesive layer can be the	30
•	30	discontinuous layer. Suitable discontinuous extensible layers include apertured non woven	
;	35	Preferred extensible apertured non woven fabrics include those made from bonded viscose filaments. Suitable fabrics of this kind are available as Bemliese (Trade Mark) from Asahai Chemical Company. Bemliese is available in weights per square metre ranging from 18 g to 45 g. A favoured fabric is Bemliese G204 which has a weight per square metre of 18.5 g.  The conformable hydrophilic polymer foam absorbent layer used in the dressing of this	35
	40	invention is adapted to be capable of absorbing the wound exudate e.g. from a burn. It is desirable that the hydrophilic foam layer absorbs the wound exudate rapidly as this enhances the low adherency properties of the dressing. Such rapid absorption prevents pooling of exudate between the dressing and the wound and it has been found that this prevention of pooling is	40
4	45	desirable.  Suitable conformable hydrophilic foams will normally be flexible, open cell foams.  The ability of open cell foams to absorb and retain fluids depends to some extent on the size	45
ţ	50	Suitable open cell hydrophilic foams of dressings of the invention have a cell size of of 30 microns to 700 microns and preferably a cell size of 50 microns to 500 microns. Apt open cell hydrophilic foams of dressings of the invention have 20% to 70% and preferably 30% to 60% of the total membrane area of the cells as membrane openings. Such open cell foams permit transport of fluid and cellular debris into and within the foam.	<b>50</b>
	55	Apt foams may be polyurethane, carboxylated butadiene styrene rubber, polyacrylate of the like foam. Such foams may be made of hydrophilic materials per se or may be treated to render them hydrophilic, or example with surfactants. It is much preferred to use foams which are	55
		can all with wound to be maintained in a moist condition even when the exudate produced has been absorbed and removed from the wind surface.	00
(	60	hydrophilic isocyanate terminated polyether prepolymer with water. Favoured hydrophilic polyurethane foams of this type include those known as Hypol foams. Hypol foams can be made polyurethane foams of this type include those known as Hypol foams. Hypol foams can be made	60
. (	65	Suitable hydrophilic foam absorbent layers have a thickness of 0.5 mm to 20 mm, more suitably 0.8 mm to 15 mm and preferably 1 mm to 12 mm.	65

	The wound dr ssings of the invention preferably consist of a conformable elastomeric apertured film, an intermediate conformable hydrophilic polymer foam layer and a continuous moisture vapour transmitting conformable polymer film outer layer in which the layers are attached in a contiguous and co-extensive relationship.	
5		5
10	It is desirable that the wound dressing of this invention are sterile. The wound dressing of the invention is advantageously provided in bacteria impervious pouches. Such packed forms can be prepared under aseptic conditions or alternatively sterilised after packing by a conventional procedure. A favoured sterilisation procedure is heat sterilisation, for example by steam. Another favoured procedure is ethylene oxide sterilisation or gamma irradiation.	
15	In another aspect the invention provides a process of making a low adherency wound dressing which comprises bringing together of a conformable elastomeric apertured film layer, an intermediate absorbent layer comprising a conformable hydrophilic polymer foam and an outer layer comprising a continuous moisture vapour transmitting conformable film.  Normally the bringing together of the layers will be a lamination process.	15
20	The previously formed individual layers can be formed into a laminate by bonding the layers together in one or more laminating processes. Suitable bonding methods include heat sealing or adhesive bonding providing the adhesive layer is moisture vapour transmitting.  In a preferred process the foam layer is formed in contact with one or both of the other layers. This process is favoured as it reduces or eliminates the number of special bonding operations.	20
25	In another preferred process the outer conformable film layer is formed on the foam layer for example by spraying a solution of the polymer.  In a continuous process the wound dressing can be made in the form of a continuous strip which is then cut up into suitable sized dressings.  The conformable hydrophilic polyurethane foam can be made by mixing together an	25
30	isocyanate terminated polyether having functionality of more than two with a surfactant and water and casting the mixture onto a surface. This surface advantageously may be the outer film of the dressing or the wound facing layer of the dressing. Preferred isocyanate terminated polyethers include Hypols FHP 2000, 2001, 3000, 3001, 2002 and 2000HD marketed by W. R. Grace & Co. Hypols are described in a booklet published by W. R. Grace and Co. "Hypol:	30
35	foamable hydrophilic polymers—laboratory procedures and foam formulation". Their preparation and use are disclosed in British Patent Specifications No. 1,429,711 and 1,507,232.  Suitable surfactants for forming conformable hydrophilic polymer foams include non-ionic surfactants. Favoured non-ionic surfactants are oxypropylene—oxyethylene block copolymers known as Pluronics marketed by BASF Wyandotte. Preferred Pluronics include L64, F87, P38,	35
40	P75 and L62. Another favoured non-ionic surfactant is a polyoxyethylene stearyl ether known as Brij 72 marketed by Honeywell Atlas.  To prepare a suitable foam 100 parts by weight of Hypol FHP 2000, 2001, 3000, 3001, 2002 or 2000HD is mixed with 0.3 to 7 parts by weight of surfactant or mixtures of surfactants	40
45	and 30 to 300 parts by weight of water and the foaming mixture cast onto a surface. Typical foaming mixtures have a cream time of about 20 secs., a rise time of about 20 secs. and a cure time of about 400 secs.  In a continuous process for forming the foam the ingredients are fed into a continuous mixing and dispensing machine. Suitable conformable hydrophilic polymer foam layers can be made by	45
50	casting the foaming mixture before it sets onto a suitable surface by means of a casting head.  A suitable mixing and dispensing machine is known as Vario-mix supplied by Prodef Engineering Limited. The foam mix can conveniently be delivered to the casting head by means of a 'fish tail' die.	50
55	In a preferred process of forming the dressing in which the foam layer is produced in contact with an external layer it is important that the other external layer should be laminated to the expanded foam while the foam is still tacky so as to obtain a good bond. Typically 2.5 minutes to 5 minutes, for example 3 mins to 3.5 mins, after the foam has been cast is suitable for bringing the foam into contact with the other external layer.	55
60	Figure 1 illustrates a process of making the conformable integral polymer nets of the wound dressing of the invention.  Figure 2 is a plan view of an embossed pattern sheet casting surface for forming an integral polymer net.  Figure 3 is a processestion through line And of Fig. 2	60
65	Figure 3 is a cross-section through line A-A of Fig. 2.  In Fig. 1 a thermoplastic film (1) with an embossed pattern on its upper surface may be fed from roll (2) to the coating head (3) where a solution (4) may be cast into the recesses of the embossed sheet. The wet cast net (5) on the embossed sheet may be passed into an oven (6) where it is dried. The dried cast net (7) may then be separated from the embossed sheet (1) and	65

wound up onto roller (8) where it may also be int rleaved with a release paper (9) fed from the roll (10). In an alternative and preferred proc ss the dried cast net is left on the embossed sheet. The coating h ad (n t shown) has an adjustable doctor blade supported on a flat bed to meter 5 the casting solution and side guides to regulate the width of the cast net. It is preferred that the doctor blade has a base portion which is thick enough to span the discrete raised areas of the embossed film to prevent the doctor blade catching in the recessed areas of the film. The doctor blade and the guides can be coated or made of a fluorocarbon polymer for example polytetrafluoroethylene to reduce friction against the film. Alternative coating heads using fixed 10 10 or rotating rollers can also be used. A favoured coating head comprises an adjustable doctor blade which is supported on a soft base, for example a base consisting of a movable rubber belt around two rotatable rollers, to meter the casting solution. Fig. 2 shows discrete raised areas (11) arranged in a square pattern to give a square grid 15 15 pattern of recesses (12) on the embossed casting sheet. In Fig. 3 a section through line A-A of Fig. 2 shows the discrete raised areas (11) in the shape of truncated square pyramids and recesses (12). The integral nets of polyurethane can be made by casting the polyurethane in a flowable state onto a surface having a pattern of discrete raised areas and interconnected recessed areas and 20 treating the case net to form a solid integral net. The flowable state of the polyurethane can 20 include solutions, dispersions, hot melts and powders which can be dried, coated, fused or otherwise to form a solid net. The casting surface may be in the form of a roller, and endless flexible belt or a length of sheet material. It is preferred that the casting surface has release properties to enable the formed net to be removed from the casting surface. The pattern of the 25 discrete raised areas and interconnected recessed areas on the casting surface selected dictates 25 the structure of the resulting net. A preferred method of making the integral nets of polyurethane is by casting a solution of a thermoplastic polyurethane onto a melt embossed polyolefin sheet and drying the cast net in a hot oven. Suitable casting solutions can contain 15% to 35% by weight of thermoplastic polyurethane, 30 preferably 20% to 30% by weight. Favoured casting solutions contain 20 to 25% by weight of Estane 5702 or Estane 5703 in acetone. Another favoured solution contains 25% to 30% by wight of Estane 5714F in tetrahydrofuran or mixtures of tetrahydrofuran and acetone. Analogous procedures may be used to prepare nets from other elastomers. The melt embossed polyolefin sheet can be made by the method given in British Patent 35 35 Specification No. 1,055,963 A suitable embossed polyolefin sheet has a pattern of 8 per cm raised areas in the form of square truncated pyramids 1 mm. wide and 0.5 mm high with side sloping to a 60°C conical angle and longitudinal and transverse square grid recesses 0.25 mm wide at the base and 0.75 mm at the top. A favoured embossed polyolefin sheet has a pattern of 6 per cm raised areas in diagonal rows 40 (45°) of square truncated pyramids 1.35 mm wide at their base, 0.7 mm wide at their top and 0.45 mm high with sides sloping to a 70° conical angle. A preferred embossed polyolefin sheet has a pattern of 4 per cm raised areas in diagonal rows (45°) of square truncated pyramids 2 mm wide at their base, 1.425 mm wide at their top and 45 45 0.5 mm high with sides sloping to a 60° conical angle. The solution of thermoplastic polyurethane can be cast onto the embossed polyolefin surface by means of a casting head consisting of a knife over a flat bed, or knife over a roller or knife over soft bed. Suitable conformable polyurethane films for the outer layer of the dressing can be formed by 50 casting or spraying from solution, hot melt coating or film extrusion in a conventional manner. 50 Preformed polyurethane films can be formed on a release surface for example a silicone release coated paper. Suitable casting solutions are described hereinbefore in relation to making integral polyurethane nets wound facing layers. In an alternative process for conformable polyurethane film can be formed on the conformable 55 hydrophilic foam layer. In an apt process the polyurethane solution is sprayed onto the foam layer. The spraying of the polyurethan solution can be carried in a conventional manner for example using an air spray gun. A suitable spray gun is model 630 available from Binks Bullow Limited. Suitable methods of making voided polyurethane-incompatible blend film outer layers are 60 60 disclosed in United Kingd m Application No. 8124250. The adhesive lay r of the outer layer can be formed by any convenient m thod including solution and emulsion coating, coating from a hot melt and by extrusion. The adhesive layer can be coated directly onto a substrate of the dressing for example the xtensible apertured non

65 woven fabric layer. However, it is preferred that the adhesive layer is coat d nto a release

surfac and the dried adhesive transferred by lamination onto the desired substrate. The wound dressing of the invention can contain topically effective medicament. Most suitably the m dicament is an antibacterial agent. Pr ferably the antibact rial agent is a broad spectrum antibacterial agent such as a silver salt such as silver sulphadiazine, an acceptable 5 iodine source such as povidone iodine (also called polyvinyl pyrrolidone iodine or PVP/I), 5 chlorhexidine salts such as the gluconate, acetate, hydrochloride or the like salts or quaternary antibacterial agents such as benzalkonium chloride or the like. A preferred medicament for inclusion in the dressing of this invention is silver sulphadiazine. A further preferred medicament for inclusion in the dressing of this invention is chlorhexidine 10 which will normally be present as one of its aforementioned salts, 10 The medicament may be present by 0.2% to 20%, more usually from 0.3 to 10% and preferably 0.5 to 5% by weight of the dressing, for example 1%, 1.2% or 3% and the like. The medicament is present in the invention in the foam layer. It is one of the surprising features of this invention that antibacterial agents can be 15 incorporated into a hydrophilic polyurethane foam and will thereafter be available to aid in 15 maintaining the wound to which the dressing is applied free of infection. It is particularly surprising that medicaments such as silver sulphadiazine and chlorhexidine hydrochloride and the like can be incorporated into the proto foam prior to polymerisation since the presence of compounds containing basic nitrogen atoms may well have been expected to 20 radically change the nature of the foam which has now been found not to occur. 20 The medicament may be introduced into the foam either by incorporation prior to foaming or by incorporation into the intact foam which has previously been prepared. If the medicament is to be introduced prior to foaming then the medicament must either be free of reactive moities which would react with the components of the mixture to be foamed (for 25 example it must not contain free amino groups which could react with the isocynates present) or 25 else the medicament must be of low solubility so that its potential reactivity is suppressed. Thus for example medicaments such as silver sulphadiazine and chlorhexidine hydrochloride are easily incorporated into the foam by dispersing the desired amount of the meidcament into the prepolymer mixture, for example dispersing it within the aqueous solution of the surfactant 30 before mixing with the isocyanate containing materials. Most suitably the insoluble medicaments 30 are in finely divided form and are most preferably micronised. It has been found that more soluble salts such as chlorhexidine gluconate cannot be incorporated in this fashion since reaction with prepolymer components can occur and a more rigid and antibacterially ineffective foam results. Fortunately it has now been discovered that 35 soluble medicaments can be included into the foam after it has been prepared by soaking the 35 foam in a solution of the medicament. Thus for example a 2 × 2 cm dressing of this invention suspended in 50mls of 5% w/v solution of chlorhexidine gluconate for 48 hours and dried was found to posess antibacterial properties. In a different embodiment of the invention the wound dressing may be in the form of a 40 compression bandage. 40 Most aptly the compression bandage of this invention has more than one elastic layer. In another aspect this invention provides a low adherency compression bandage which comprises a wound facing layer of elastomeric apertured film, an intermediate layer of a conformable hydrophilic foam and an outer layer of an elastomeric continuous moisture vapour permeable 45 film. 45 Most aptly the elastomeric apertured film is a net as hereinbefore described. In preferred compression bandages of this invention the integral net wound facing layer, the intermediate foam layer and the outer film layer are made of polyurethane as hereinbefore described. Low adherency compression bandages of the invention can be used to cover skin grafts. In 50 this application the bandage may absorb exudate from the graft surrounds and at the same time exert an even pressure over the graft site which can protect against hypertrophic scarring. The low adherency compression bandages of this invention may also be used in the treatment of ulcers, for example leg ulcers. The bandages of this invention will normally be presented as rolls of from 1 to 4 meters 55 length and 5 to 20cms width. After fabrication the wound dressing of this invention can be washed with water to remove excess surfactant and then dried. Surprisingly the method of drying has been found to greatly affect the appearance of th 60 Some air dried dressings have been found to curl. In some cases the outer film has puckered th reby giving the film a grained appearanc and texture of skin. Dressings autoclaved using vacuum drying cycle have been f und to tend to remain flat. The outer film layer has a puckered surface. Partially dried dressings, that is dressings containing 65 low levels of residual water absorbed into the foam polymer but not into foam air spaced have 65

		been discovered to be flat. This unanticipated ffect is rendered ven more useful since the dressing will remain flat if protected against loss of water, for example if packaged in a water proof pouch such as an aluminium foil pouch. In such partially dried dressings the film and net are not puckered.	٠.
	5		5
	10	As previously indicated hereinbefore the dressings of this invention may be adapted to release an antibacterially effective amount of an antibacterial agent into the wound covered by the dressing. Thus in an alternative aspect this invention provides a method of treating a wound so as to aid in rendering or maintaining it free of infection which comprises contacting the wound with a dressing of this invention adapted to release an antibacterial agent. Most aptly this aspect of the invention is employed in rendering or maintaining burns free of infection. The	10
	15	antibacterial agent present is favourably a silver salt such as silver sulphadiazine or a chlorhexidine salt such as chlorhexidine hydrochloride or a mixture thereof. Preferably the antibacterial agent present is silver sulphadiazine.  The absence of fibres in the dressing enhances the non adherent properties of the wound dressings of the invention.	15
	20	Description Preparation of Integral Polyurethane Net A solution containing 30% by weight of Estane 5714F1 in tetrahydrofuran was cast into the recesses of a 15cm wide melt embossed high density polyethylene sheet by means of the blade	20
	25	over flat bed spreading technique. The sheet had a melt embossed pattern of 8 per cm raised areas in the form of square truncated pyramids 1mm wide at their base and 0.5mm high with sides sloping to a solid conical angle of 60°C. The wet cast net on the embossed film was dried by passage through a hot air circulating oven at a temperature of 90°C to 100°C for two minutes. The dried cast net was separated from the embossed film and wound onto a roller interleaved with a double sided silicone release paper.	25
	30	The resultant cast integral of elastomeric polyurethane net had the following properties: Weight 40gsm; thickness $100-125$ microns; aperture size 0.3 to 0.4mm; tensile strength (g/2.5cm wide), machine direction $800 \pm 51$ , transverse direction $664 \pm 57$ , elongation at break %, machine direction $389 \pm 24$ , transverse direction $374 \pm 24$ .	30
	35	Preparation of Integral Polyurethane Diamond Pattern Net (6 apertures/cm).  The net was cast in the same manner as the preceeding square net except that the casting sheet had a melt embossed pattern of 6 per cm raised areas in diagonal rows (45°) of square truncated pyramids 1.35mm wide at their base and 0.45mm high with sides sloping to a conical angle of 70°	35
	40	Example 1 Preparation of a low adherency wound dressing	40
	45	A mixture of Brij 72 (22.5g of 2% aqueous emulsion) and Pluronic F87 (0.5ml of a 10% aqueous solution) was added to Hypol FHP3001 (15g) in a beaker and thoroughly mixed by stirrriing with a metal spatula until the Hypol ws uniformly dispersed (20 seconds). The foaming mixture was poured into a 15cm wide brass hand spreader box set at a gap of 1.8mm above a 25 micron thick cast Estane 5714F1 film. The spreader box was then drawn by hand along the film surface to leave a foam layer on the film The cast integral polyurethane net of the	45
	50	description was then laminated to the foam by placing the net (smooth surface uppermost) onto the setting foam 3 minutes to 3.25 minutes after the spreading had commenced. The foam layer was free of large craters and was well bonded to the net.  Alternatively the wound dressing can be made in a similar manner by coating the foam onto the integral polyurethane net (preferably on the embossed film carrier) and laminating the film to	50
-	55	the setting foam.  Sample wound dressings of Example 1 and the comparison hydrophilic polyurethane foam were washed with distilled water and dried at 40°C for 12 hours before being tested for wound adherency.	55
	60	Example 2 Brij 72 (30g as a 2.5% aqueous solution) was added to Hypol FHP 3001 (20 g) in a beaker and mixed by stirring with a metal spatula and then with a mechanical stirrer until the Hypol was uniformly dispersed (approximately 20 seconds). The foaming mixture was cast onto a 6 aperture/cm diamond pattern net as described above n its emboss d film carrier by m ans of a blade over flat bed coating head set at a gap of 0.1 mm. A 25 micron thick film (Estan 5714F1) on a silicone release paper was laminated to th foam surface with light manual pressure to the foam surface while it was still tacky (approximately 3 minutes after spreading).	60
	-		

	After 15 minut s the mboss d film and silicone coated release papers were removed from the n t and film surfaces of the strip and the strip cut int $30 \text{ cm} \times 15 \text{ cm}$ wound dressings. The dressings were washed in two changes of 1 litre of distilled water and dried in air.	
5	Example 3 Wound dressings were prepared in the same manner as Example 2 using a gap setting of 0.5 mm instead of 0.1 mm.	5
10	Example 4 Wound dressings were prepared in the same manner as Example 2 using a gap setting of 1.0 mm instead of 0.1 mm.	10
15	Example 5 The wound dressings of Example 3 were autoclaved at 116°C for 30 minutes followed by a vacuum drying cycle.	15
	Example 6 The wound dressings of Example 4 were autoclaved in the same manner as Example 5.	
20	Example 7 The dressings of Example 4 were washed but only partially dried by padding with an absorbent towel.	20
25	The following Examples 8 to 12 the preparation of of medicated wound dressings of the invention.	25
30	Example 8 Wound dressings were prepared in the same manner as Example 3 except that silver sulphadiazine powder (0.2g) was blended into the Brij 72 emulsion with a high speed shear mixer prior to the addition of Hypol FHP 3001.	30
35	Example 9 Wound dressings were prepared in the same manner as Example 8 using 1g of chlorhexidine hydrochloride powder instead of silver sulphadiazine powder (0.2g).	35
	Example 10 Wound dressings prepared as in Example 3 were soaked for 10 minutes in a tray containing an aqueous solution of chlorhexidine gluconate (5% weight/volume) and air dried.	
40	Example 11 Wound dressings were prepared in the same manner as Example 10 using an aqueous solution of chlorhexidine acetate (5% weight/volume) instead of an aqueous solution of chlorhexidine gluconate.	40
45	Example 12 Wound dressings were prepared in the same manner as Example 10 using an aqueous solution of povidone iodine (10% weight/volume) instead of an aqueous solution of chlorhexidine gluconate.	45
50	Example 13 Wound dressings were prepared in the same manner as Example 7 using 20 g instead of 30 g of Brij 72.	50
55	Example 14 Wound dressings were prepared in the same manner as Example 7 using 40 g instead of 30 g of Brij 72.	55
60	Example 15 Wound dressings were prepared in the sam manner as Example 7 with 0.5 ml of Pluronic L64 (10% aqueous solution) added to the surfactant mulsion.	60
00	Example 16 Wound dressings were prepared in the same manner as Example 15 using Pluronic F68 in place of Pluronic L64.	- <b>-</b>
65	Example 17	65

Wound dressings were prepared in the same manner as Example 15 using Pluronic F108 instead of Pluronic L64.

Example 18

Wound dressings were prepared in the same manner as Example 1 using Brij 72 (30g as a 2.5% aqueous emulsion) instead of a mixture of Brij 72 (22.5g as a 2% aqueous emulsion) and Pluronic F87 as a 10% aqueous solution).

Example 19

Wound dressings were prepared as Example 18 using a mixture of Brij 72 and a Pluronic P75 10 (0.5 ml as a 10% aqueous solution).

Example 20

Wound dressings were prepared in the same manner as Example 1 using a cast polybutadiene 15 (ref RB830) net (8 apertures per cm) instead of a polyurethane net. 15

Example 21

Preparation of a Compression Bandage

A bandage strip was made in the same manner as the wound dressing strip of Example 2 20 using a blade gap setting of 0.5 mm. The strip was washed in distilled water and dried in air. A 20 1 cm wide bandage had the following stress-strain properties:

Strain %	Stress g/cm
25	360
50	610
100	800
	50

25

To illustrate the compression properties of the bandage the pressure (P) exerted on a limb of radius (R) by a single layer of the bandage stretched by different amounts was calculated from the stress-strain properties above using the equation

35 p = 
$$\frac{F}{P}$$

35

30

where F is the force (g/cm) to stretch the bandage by a given amount.

40	0				_	40
	Arm 10 cm in diameter	stretched length (% of original length)	g/cm <sup>-2</sup>	<sup>P</sup> mmHg		
45		125 150	72 122	54 92		45
		200	160	120		
	Leg 20cm in	125	36	27		50
50	diameter	150 200	61 80	46 60		

Antibacterial properties of the Medicated Dressings

15 mm diameter samples of medicated wound dressings of Examples 9 to 14 were moistened 55 and placed on agar plates seeded with staphyloccocus aureus or bacillus subtilis and incubated for 24 hours at optimum t mperature for growth. The zones of inhibition produced by diffusion of th medicament from the dressings were measured.

						•		
	Example	Medicament Zone diameter						
5				Staph	.B. subtilis	,	5	
				aureus				
	8	silver sulphadjazine		30	29.5			
	9	chlorhexidine Hcl	•	23.8	21.6			
	12	povidone iodine		29.6	19.4		10	
10		chlorhexidine acetate chlorhexidine gluconate	/1\	29.0 22.6	32.0 23.5		10	
	10*	chiornexidine glucoriate	(1) (2)	19.8	19.1			
			(3)	16.8	16.0			
	control (no	medicament)	(0)	0	0			
15	*consecu	tive tests on the same samp	le to me	easure the	effect of the		15	
	sustained r	elease of the medicament fr	om the	dressing.				
-00	5	•					20	
20	Example 2	2 n of Integral Diamond Patter	n Not				20	
	(4 aperture		11 1101					
	A solutio	n containing 20% by weigh	t of Esta	ane 5714F	in 60/40 (wei	ght by weight) mixture		
	of Tetrahyo	Irofuran /acetone was cast i	nto the	recesses of	f a 15 cm wide	melt embossed high		
25	density not	vethylene sheet by means of	f a blade	e over soft	bed coating ted	chnique. The sheet had	25	
	a melt emb	ossed pattern of 4 per cm r	aised an	eas in diag	onal rows (45°)	of square truncated		
	pyramids 2	mm wide at their base, 1.4 I angle of 60°. The wet cast	2 mm v	vide at top	and U.5 mm n	ad by passage through		
	to a conica	ren at temperature of 80°C	for two	nio embos: minutes	eu IIIII was un	ed by passage timough		
30	The net	had a weight per square me	tre of 33	3 g and ha	d 4 per cm ape	rtures of approximately	30	
30	1.4 mm in			<b>.</b>	<b>—</b> • <b>F</b> • • • • • • • • • • • • • • • • • • •	,		
	Preparation	of the absorbent layer			.P. Alb. Dade	Fundamentary I tradepolity		
~=	Using a two component dispensing unit (Vario-mix supplied by Prodef Engineering Limited), a foaming mixture was formed by mixing Hypol FHP 2002 and Brij 72 (2% aqueous solution) in						35	
35	the ratio of 1:2.25. The foaming mixture was fed into the coating head by means of an output							
	nozzle in th	ne form of a 15 cm 'fish tail	die' and	i coated or	nto the cast poly	yurethane net (on		
	embossed :	film) by means of a knife over	er roller	coating he	ead set at a gap	of 1 mm. The cast		
foam was dried by passage through an air circulating oven at a temperature of 50°C for 5								
40	40 minutes.							
	Preparation of the Conformable Film Outer Layer A solution containing 2% by weight a polyurethane of (Estane 5714F) in a 60/40 (weight I							
	weight) mixture of tetrahydrofuran/acetone was hand sprayed onto the foam surface of the							
45	composite foam/net strip using an air spray unit (model 630 supplied by Binks Bullow Limited)							
40	and dried by passage through an air circulating oven heated to a temperature of 70°C.							
	The polyurethane coating was found to be continuous and had a weight per square metre of							
	approximat	ely 30 grams.				atrin and the atrin and		
<b>E</b> 0	The emb	ossed film was then remove	a trom t	ine three is	iyer composite :	strip and the strip cut	50	
50	into aressir	ngs of suitable size for adher	ency les	sung.			•	
	Example 2	<i>3</i>						
	A foam-r	net composite strip on embos	ssed filn	n was mad	e in the same n	nanner as example 22.		
	An exten	sible apertured non woven f	abric (B	emliese GS	3 204) coated v	vith a continuous layer	55	
55	of a moistu	re vapour transmitting acryl	ylate copolymer adhesive (30 g/m²) was laminated to the					
	foam side	of the composite strip to form	n a cont	rormable o	uter layer.	47 parts by waight of		
	The adhesive was a copolymer of 47 parts by weight n-butyl acrylate, 47 parts by weight of							
	2-ethyl-hexyl acrylate and 6 parts by weight of acrylic acid having an intrinsic viscosity of 1.9 dl/g polymerised in aceton according t the method given in United Kingdom Application No.							
60	8106707							
50	The mb	mbossed film was then removed fr m th composite strip and th strip cut into suitable						
		ings of the invention.	•	•	-			
<u></u>	Example 2	4	L		a Evampla 22 ··	icina a cast polyhistadi.	65	
65	vvound	dressings were prepared in the	ne sam	manner a	s example 22 U	ionina a cast honanarani.	00	
	·							

ene (Ref RB 830) net (6 apertures/cm) instead of a polyurethane net.

Adherency Testing When tested on guinea pigs the dressing of Examples 4, 24 and 26 required only half the energy to remove them than did "Melolin" (a commercial non-adherent dressing) and Examples 5 1 and 2 required respectively about one eighth and one third the energy to remove them than did "Melolin". **CLAIMS** 1. A low adherency wound dressing which comprises a wound facing layer, an intermediate 10 absorbent layer and an outer layer which wound dressing is characterised in that the wound facing layer comprises a conformable elastometic apertured film, the intermediate absorbent layer comprises a conformable hydrophilic foam and the outer layer comprises a continuous moisture vapour transmitting conformable film. 2. A low adherency wound dressing as claimed in claim 1 in which the conformable 15 elastomeric apertured film comprises an integral net. 3. A low adherency wound dressing as claimed in either of claims 1 or 2 in which the conformable elastomeric apertured film has 4 to 40 apertures per cm with a dimension 0.05 mm to 2.5mm. 4. A low adherency wound dressing as claimed in any of claims 1 to 3 in which the 20 conformable elastomeric apertured film comprises a polyurethane. 5. A low adherency wound dressing as claimed in any of claims 1 to 4 in which the continuous conformable film of the outer layer has a moisture vapour transmission of 500 to 2000 grams/square metre/24 hours at 37.5°C at 100% to 10% relative humidity difference. 6. A low adherency wound dressing as claimed in any of claims 1 to 5 in which the 25 continuous moisture vapour transmitting conformable film comprises a polyurethane. 7. A low adherency wound dressing as claimed in any of claims 1 to 8 in which the conformable film outer layer comprises a moisture vapour transmitting adhesive layer. 8. A low adherency wound dressing as claimed in any of claims 1 to 7 in which the 30 30 conformable hydrophilic foam is a foam of a hydrophilic polymer. 9. A low adherency wound dressing as claimed in claim 8 in which the hydrophilic polymer is a hydrophilic polyurethane. 10. A low adherency wound dressing as claimed in any of claims 1 to 9 in which the hydrophilic foam comprises an open cell foam with a cell size of 50 microns to 500 microns. 11. A low adherency wound dressing as claimed in any of claims 1 to 10 in which the 35 hydrophilic foam comprises an open cell foam in which 30% to 60% of the total membrane area of the cells are membrane openings. 12. A low adherency wound dressing as claimed in any of claims 1 to 11 in which the dressing contains a topically effective medicament. 13. A sterile low adherency wound dressing as defined in claims 1 to 12 within a bacteria 40

14. A low adherency compression bandage which comprises a wound facing layer of elastomeric apertured film, an intermediate layer of conformable hydrophilic foam and an outer

layer of elastomeric continuous moisture vapour permeable film.

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impervious pack.